# MPICH2 User's Guide\* Version 0.4 Mathematics and Computer Science Division Argonne National Laboratory

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### 1 Introduction

This manual assumes that MPICH2 has already been installed. For instructions on how to install MPICH2, see the MPICH2 Installer's Guide, or the README in the top-level MPICH2 directory. This manual explains how to compile, link, and run MPI applications, and use certain tools that come with MPICH2. This is a preliminary version and some sections are not complete yet. However, there should be enough here to get you started with MPICH2.

## 2 Setting Paths

You will have to know the directory where MPICH2 has been installed. (Either you installed it there yourself, or your systems administrator has installed it. One place to look in this case might be <code>/usr/local.</code>) We suggest that you put the <code>bin</code> subdirectory of that directory in your path. This will give you access to assorted MPICH2 commands to compile, link, and run your programs conveniently. Other commands in this directory manage parts of the run-time environment and execute tools.

One of the first commands you might run is mpich2version to find out the exact version and configuration of MPICH2 you are working with. Some of the material in this manual depends on just what version of MPICH2 you are using and how it was configured at installation time.

# 3 Quick Start

You should now be able to run an MPI program. Let us assume that the directory where MPICH2 has been installed is /home/you/mpich2-installed, so that in the section above you did

setenv PATH /home/you/mpich2-installed/bin: \$PATH

for tcsh and csh, or

expoert PATH=/home/you/mpich2-installed/bin:\$PATH

for bash or sh. Then to run an MPI program, albeit only on one machine, you can do:

```
mpd &
cd /home/you/mpich2-installed/examples
mpiexec -n 3 cpi
mpdallexit
```

Details for these commands are provided below, but if you can successfully execute them here, then you have a correctly installed MPICH2 and have run an MPI program.

# 4 Compiling and Linking

A convenient way to compile and link your program is by using scripts that use the same compiler that MPICH2 was built with. These are mpicc, mpicxx, mpif77, and mpif90, for C, C++, Fortran 77, and Fortran 90 programs, respectively. If any of these commands are missing, it means that MPICH2 was configured without support for that particular language.

## 4.1 Specifying Compilers

You need not use the same compiler that MPICH2 was built with, but not all compilers are compatible. You can also specify the compiler for building MPICH2 itself, as reported by mpich2version just by using the compiling and linking commands from the previous section. (See the *Installer's Guide*).

#### 4.2 Shared Libraries

Currently shared libraries are only tested on Linux, and there are restrictions. See the *Installer's Guide* for how to build MPICH2 as a shared library.

### 4.3 Special Issues for Fortran

This section is under development.

- Review of basic and extended support; the Fortran 90 module.
- Various name-mangling issues

# 5 Running Programs with mpiexec

If you have been using the original MPICH, or any of a number of other MPI implementations, then you have probably been using mpirun as a way to start your MPI programs. The MPI-2 Standard describes mpiexec as a suggested way to run MPI programs. MPICH2 implements the mpiexec standard, and also provides some extensions. MPICH2 provides mpirun for backward compatibility with existing scripts, but it does not support the same or as many options as mpiexec.

## 5.1 Standard mpiexec

Here we describe the standard mpiexec arguments from the MPI-2 Standard [1]. The simplest form of a command to start an MPI job is

```
mpiexec -n 32 a.out
```

to start the executable a.out with 32 processes (providing an MPI\_COMM\_WORLD of size 32 inside the MPI application). Other options are supported, for specifying hosts to run on, search paths for executables, working directories, and even a more general way of specifying a number of processes. Multiple sets of processes can be run with different exectuables and different values for their arguments, with ":" separating the sets of processes, as in:

```
mpiexec -n 1 -host loginnode master : -n 32 -host smp slave
```

The -configfile argument allows one to specify a file containing the specifications for process sets on separate lines in the file. This makes it unnecessary to have long command lines for mpiexec. (See p. 353 of [2].)

Currently the -soft argument (for giving hints instead of a precise number for the number of processes) is not supported.

It is also possible to start a one process MPI job (with size of MPI\_COMM\_WORLD equal to 1), without using mpiexec. This process will become an MPI process when it calls MPI\_Init, and can then call other MPI functions, including MPI\_Comm\_spawn.

#### 5.2 Extensions for All Process Management Environments

Some mpiexec arguments are specific to particular communication subsystems ("devices") or process management environments ("process managers"). Our intention is to make all arguments as uniform as possible across devices and process managers. For the time being we will document these separately.

# 5.3 Extensions for the MPD Process Management Environment

MPICH2 provides a number of process management systems. The default is called MPD. MPD provides a number of extensions to the standard form of mpiexec.

#### 5.3.1 mpiexec arguments for MPD

The default configuration of MPICH2 chooses the MPD process manager and the "simple" implementation of the Process Management Interface. MPD provides a version of mpiexec that supports both the standard arguments described in Section 5.1 and other arguments described in this section. MPD also provides a number of commands for querying the MPD process management environment and interacting with jobs it has started.

Before running mpiexec, the runtime environment must be established. In the case of MPD, the daemons must be running. See Section 6.1 for how to run and manage the MPD daemons.

We assume that the MPD ring is up and the installation's bin directory is in your path; that is, you can do:

mpdtrace

and it will output a list of nodes on which you can run MPI programs. Now you are ready to run a program with mpiexec. Let us assume that you have compiled and linked the program cpi (in the installdir/examples directory and that this directory is in your PATH. Or that is your current working directory and '.' ("dot") is in your PATH. The simplest thing to do is

```
mpiexec -n 5 cpi
```

to run cpi on five nodes. The process management system (such as MPD) will choose machines to run them on, and cpi will tell you where each is running.

You can use mpiexec to run non-MPI programs as well. This is sometimes useful in making sure all the machines are up and ready for use. Useful examples include

```
mpiexec -n 10 hostname
```

and

mpiexec -n 10 printenv

#### 5.3.2 Passing Environment Variables to Processes

The MPI-2 standard specifies the syntax and semantics of the arguments -n, -path,-wdir, -host, -file, -configfile, and -soft. All of these are currently implemented for MPD's mpiexec except -soft. Each of these is what we call a "local" option, since its scope is the processes in the set of processes described between colons, or on separate lines of the file specified by -configfile. We add some extensions that are local in this way and some that are "global" in the sense that they apply to all the processes being started by the invocation of mpiexec.

The MPI-2 Standard provides a way to pass different arguments to different application processes, but does not provide a way to pass environment variables. The local parameter <code>-env</code> does this for one set of processes. That is,

```
mpiexec -n 1 -env FOO BAR a.out : -n 2 -env BAZZ FAZZ b.out
```

makes BAR the value of environment variable FOO on the first process, running the executable a.out, and gives the environment variable BAZZ the value FAZZ on the second two processes, running the executable b.out. To set an environment variable without giving it a value, use '' as the value in the above command line.

The global parameter **-genv** can be used to pass the same environment variables to all processes. That is,

```
mpiexec -genv FOO BAR -n 2 a.out : -n 4 b.out
```

makes BAR the value of the environment variable F00 on all six processes. If -genv appears, it must appear in the first group. If both -genv and -env are used, the -env's add to the environment specified or added to by the -genv variables. If there is only one set of processes (no ":"), the -genv and -env are equivalent.

The local parameter -envall is an abbreviation for passing the entire environment in which mpiexec is executed. The global version of it is -genvall. This global version is implicitly present. To pass no environment variables, use -envnone and -genvnone. So, for example, to set only the environment variable FOO and no others, regardless of the current environment, you would use

```
mpiexec -genvnone -env FOO BAR -n 50 a.out
```

A list of environment variable names whose values are to be copied from the current environment can be given with the -envlist (respectively, -genvlist) parameter; for example,

```
mpiexec -genvnone -envlist PATH, LD_SEARCH_PATH -n 50 a.out
```

sets the PATH and LD\_LIBRARY\_PATH in the environment of the a.out processes to their values in the environment where mpiexec is being run. In this situation you can't have commas in the environment variable names, although of course they are permitted in values.

Some extension parameters have only global versions. They are

-1 provides rank labels for lines of stdout and stderr. These are a bit obscure for processes that have been explicitly spawned, but are still useful.

- -usize sets the "universe size" that is retrieved by the MPI attribute MPI\_UNIVERSE\_SIZE on MPI\_COMM\_WORLD.
- -bnr is used when one wants to run executables that have been compiled and linked using the ch\_p4mpd or myrinet device in MPICH1. The MPD process manager provides backward compatibility in this case.

#### 5.3.3 Environment Variables Affecting mpiexec

A small number of environment variables affect the behavior of mpiexec.

- MPIEXEC\_TIMEOUT The value of this environment variable is the maximum number of seconds this job will be permitted to run. When time is up, the job is aborted.
- MPIEXEC\_BNR If this environment variable is defined (its value, if any, is currently insignificant), then MPD will act in backward-compatibility mode, supporting the BNR interface from the original MPICH (e.g. versions 1.2.0 1.2.6) instead of its native PMI interface, as a way for application processes to interact with the process management system.

#### 5.4 Extensions for SMPD

SMPD is an alternate process manager that runs on both Unix and Windows. It can launch jobs across both platforms if the binary formats match (big/little endianness and size of C types - int,long,void\*,etc).

#### 5.4.1 mpiexec arguments for SMPD

mpiexec for smpd accepts the standard MPI-2 mpiexec options. Execute

mpiexec

or

mpiexec -help2

to print the usage options. Typical usage:

```
mpiexec -n 10 myapp.exe
```

All options to mpiexec:

-n x

-np x

launch x processes

-localonly x

-np x -localonly

launch x processes on the local machine

-machinefile filename

use a file to list the names of machines to launch on

-host hostname

launch on the specified host.

-hosts n host1 host2 ... hostn

-hosts n host1 m1 host2 m2 ... hostn mn

launch on the specified hosts. In the second version the number of processes = m1 + m2 + ... + mn

-dir drive:\my\working\directory

-wdir /my/working/directory

launch processes with the specified working directory. (-dir and -wdir are equivalent)

-env var val

set environment variable before launching the processes

-exitcodes

print the process exit codes when each process exits.

-noprompt

prevent mpiexec from prompting for user credentials. Instead errors will be printed and mpiexec will exit.

-port port

#### -p port

specify the port that smpd is listening on.

#### -phrase passphrase

specify the passphrase to authenticate connections to smpd with.

#### -smpdfile filename

specify the file where the smpd options are stored including the passphrase. (unix only option)

## -soft Fortran90\_triple

acceptable number of processes to launch up to maxprocs

#### -path search\_path

search path for executable,; separated

#### -timeout seconds

timeout for the job.

#### -prompt

Windows specific options:

### -map drive:\\host\share

map a drive on all the nodes this mapping will be removed when the processes exit

#### -logon

prompt for user account and password

#### -pwdfile filename

read the account and password from the file specified put the account on the first line and the password on the second

#### -nomapping

don't try to map the current directory on the remote nodes

#### -nopopup\_debug

disable the system popup dialog if the process crashes

#### -dbg

catch unhandled exceptions

#### -priority class[:level]

set the process startup priority class and optionally level.

class = 0,1,2,3,4 = idle, below, normal, above, high

level = 0.1, 2.3, 4.5 = idle, lowest, below, normal, above, highest

the default is -priority 1:3

#### -register

encrypt a user name and password to the Windows registry.

#### -remove

delete the encrypted credentials from the Windows registry.

#### -validate [-host hostname]

validate the encrypted credentials for the current or specified host.

#### 5.5 **Extensions for Forker**

The forker is a process management system for starting processes on a single machine, so called because the MPI processes are simply forked from the mpiexec process.

#### 5.5.1mpiexec arguments for Forker

The argument -maxtime sets a maximum time in seconds for the job to run.

#### 6 Managing the Process Management Environment

Some of the process managers supply user commands that can be used to interact with the process manager and to control jobs. In this section we describe user commands that may be useful.

#### 6.1 MPD

mpd starts an mpd daemon.

mpdboot starts a set of mpd's on a list of machines.

mpdtrace lists all the MPD daemons that are running. The -1 option lists full hostnames and the port where the mpd is listening.

mpdlistjobs lists the jobs that the mpd's are running. Jobs are identified by the name of the mpd where they were submitted and a number.

mpdkilljob kills a job specified by the name returned by mpdlistjobs

mpdsigjob delivers a signal to the named job. Signals are specified by name or number.

You can use keystrokes to provide signals in the usual way, where mpiexec stands in for the entire parallel application. That is, if mpiexec is being run in a Unix shell in the foreground, you can use ^C (control-C) to send a SIGINT to the processes, or ^Z (control-Z) to suspend all of them. A suspended job can be continued in the usual way.

Precise argument formats can be obtained by passing any MPD command the --help or -h argument. More details can be found in the README in the mpich2 top-level directory or the README file in the MPD directory mpich2/src/pm/mpd.

# 7 Debugging

Debugging parallel programs is notoriously difficult. Here we describe a number of approaches, some of which depend on the exact version of MPICH2 you are using.

### 7.1 mpigdb

If you are using the MPD process manager, you can use the command mpigdb instead of mpiexec to execute a program with each process running under the control of the gdb sequential debugger. mpigdb helps control the multiple instances of gdb by sending stdin either to all processes or to a selected process and by labeling and merging output. The following script of an mpigdb session gives an idea of how this works. Input keystrokes are sent to all processes unless specifially directed by the "z" command.

```
0-9: (gdb) 1
0-9: 5 double f(double);
0-9: 6
0-9: 7 double f(double a)
0-9: 8 {
0-9: 9
           return (4.0 / (1.0 + a*a));
0-9: 10
               }
0-9: 11
0-9: 12
               int main(int argc,char *argv[])
0-9: 13
0-9: 14
                   int done = 0, n, myid, numprocs, i;
0-9: (gdb)
0-9: 15
                   double PI25DT = 3.141592653589793238462643;
0-9: 16
                   double mypi, pi, h, sum, x;
0-9: 17
                   double startwtime = 0.0, endwtime;
0-9: 18
                   int namelen:
0-9: 19
                   char processor_name[MPI_MAX_PROCESSOR_NAME];
0-9: 20
0-9: 21
                   MPI_Init(&argc,&argv);
0-9: 22
                   MPI_Comm_size(MPI_COMM_WORLD,&numprocs);
0-9: 23
                   MPI_Comm_rank(MPI_COMM_WORLD,&myid);
0-9: 24
                   MPI_Get_processor_name(processor_name,&namelen);
0-9: (gdb)
0-9: 25
0-9: 26
                   fprintf(stdout, "Process %d of %d is on %s\n",
0-9: 27
                           myid, numprocs, processor_name);
0-9: 28
                   fflush(stdout);
0-9: 29
0-9: 30
                   n = 10000;
                                               /* default # of rectangles */
0-9: 31
                   if (myid == 0)
0-9: 32
                       startwtime = MPI_Wtime();
0-9: 33
0-9: 34
                   MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
0-9: (gdb) b 30
0-9: Breakpoint 2 at 0x400000000002541: file /home/lusk/mpich2/examples/cpi.c, line 30.
0-9: (gdb) r
0-9: Continuing.
0: Process 0 of 10 is on ksl2
1: Process 1 of 10 is on ksl2
2: Process 2 of 10 is on ksl2
3: Process 3 of 10 is on ksl2
4: Process 4 of 10 is on ksl2
5: Process 5 of 10 is on ksl2
6: Process 6 of 10 is on ksl2
7: Process 7 of 10 is on ksl2
```

```
8: Process 8 of 10 is on ksl2
9: Process 9 of 10 is on ksl2
0-9:
0-9: Breakpoint 2, main (argc=1, argv=0x60000fffffffb4b8)
0-9:
         at /home/lusk/mpich2/examples/cpi.c:30
0-9: 30
                   n = 10000;
                                               /* default # of rectangles */
0-9: (gdb) n
0-9: 31
                   if (myid == 0)
0-9: (gdb) n
0: 32
               startwtime = MPI_Wtime();
1-9: 34
                   MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
0-9: (gdb) z 0
0: (gdb) n
0: 34
           MPI_Bcast(&n, 1, MPI_INT, 0, MPI_COMM_WORLD);
0: (gdb) z
0-9: (gdb) where
0-9: #0 main (argc=1, argv=0x60000fffffffb4b8)
         at /home/lusk/mpich2/examples/cpi.c:34
0-9: (gdb) n
0-9: 36
                   h = 1.0 / (double) n;
0-9: (gdb)
0-9: 37
                   sum = 0.0;
0-9: (gdb)
0-9: 39
                   for (i = myid + 1; i \le n; i += numprocs)
0-9: (gdb)
0-9: 41
                       x = h * ((double)i - 0.5);
0-9: (gdb)
0-9: 42
                       sum += f(x);
0-9: (gdb)
                   for (i = myid + 1; i \le n; i += numprocs)
0-9: 39
0-9: (gdb)
                       x = h * ((double)i - 0.5);
0-9: 41
0-9: (gdb)
0-9: 42
                       sum += f(x);
0-9: (gdb)
0-9: 39
                   for (i = myid + 1; i <= n; i += numprocs)
0-9: (gdb)
0-9: 41
                       x = h * ((double)i - 0.5);
0-9: (gdb)
0-9: 42
                       sum += f(x);
0-9: (gdb)
0-9: 39
                   for (i = myid + 1; i <= n; i += numprocs)
0-9: (gdb)
0-9: 41
                       x = h * ((double)i - 0.5);
0-9: (gdb)
```

```
0-9: 42
                      sum += f(x);
0-9: (gdb)
0-9: 39
                 for (i = myid + 1; i <= n; i += numprocs)
0-9: (gdb)
0-9: 41
                      x = h * ((double)i - 0.5);
0-9: (gdb)
0-9: 42
                      sum += f(x);
0-9: (gdb)
0-9: 39
                 for (i = myid + 1; i <= n; i += numprocs)
0-9: (gdb)
                    x = h * ((double)i - 0.5);
0-9: 41
0-9: (gdb)
0-9: 42
                     sum += f(x);
0-9: (gdb) p sum
0: $1 = 19.999875951497799
1: $1 = 19.999867551672725
2: $1 = 19.999858751863549
3: $1 = 19.999849552071328
4: $1 = 19.999839952297158
5: $1 = 19.999829952542203
6: $1 = 19.999819552807658
7: $1 = 19.999808753094769
8: $1 = 19.999797553404832
9: $1 = 19.999785953739192
0-9: (gdb) c
0-9: Continuing.
0: pi is approximately 3.1415926544231256, Error is 0.0000000008333325
1-9:
1-9: Program exited normally.
1-9: (gdb) 0: wall clock time = 44.909412
0: Program exited normally.
0: (gdb) q
0-9: MPIGDB ENDING
ks12%
```

You can attach to a running job with

```
mpdgdb -a <jobid>
```

where <jobid> comes from mpdlistjobs.

## 8 MPICH2 under Windows

#### 8.1 Directories

The default installation of MPICH2 is in C:\Program Files\MPICH2. Under the installation directory are three sub-directories: include, bin, and lib. The include and lib directories contain the header files and libraries necessary to compile MPI applications. The bin directory contains the process manager, smpd.exe, and the MPI job launcher, mpiexec.exe. The dlls that implement MPICH2 are copied to the Windows system32 directory.

## 8.2 Compiling

The libraries in the lib directory were compiled with MS Visual C++.NET 2003 and Intel Fortran 8.0 with the default MPICH2 socket channel. These compilers and any others that can link with the MS .lib files can be used to create user applications. gcc and g77 for cygwin can be used with the libmpich\*.a libraries.

For MS Developer Studio users: Create a project and add C:\Program Files\MPICH2\include to the include path and C:\Program Files\MPICH2\lib to the library path. Add cxx.lib and mpich2.lib to the Release target link command. Add cxxd.lib and mpich2d.lib to the Debug target.

Intel Fortran 8 users add fmpich2d.lib to the link command in addition to the libraries mentioned above.

cygwin users use libmpich2.a libfmpich2g.a.

#### 8.3 Running

MPI jobs are run from a command prompt using mpiexec.exe. See section 5.4 on mpiexec for smpd for a description of the options to mpiexec.

# References

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